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Facile Synthesis of a Stable Dihydroboryl $\{BH_2\}^-$ Anion

Merle Arrowsmith, James D. Mattock, Stephan Hagspiel, Ivo Krummenacher, Alfredo Vargas and Holger Braunschweig*

Abstract: While the one-electron reduction of $(CAAC^{Me})BH_2Br$ ($CAAC^{Me} = 1-(2,6-diisopropylphenyl)-3,3,5,5-tetramethylpyrrolidin-2-ylidene$) yields a hydride-shift isomer of the corresponding tetrahydrodiborane, a further reversible reduction leads to the first stable parent boryl anion, $[(CAAC^{Me})BH_2]^-$, which acts as a powerful boron nucleophile.

Tricoordinate boron reagents generally behave as electrophiles due to their vacant p orbital. In 2006, however, Yamashita and Nozaki reported the first boryl anion, compound **I** (Fig. 1),^[1] in which boron is in its formal +1 oxidation state and reacts as a nucleophile, thus enabling access to a wide range of novel boron-element-bonded compounds.^[2] Curran and coworkers later succeeded in generating a highly reactive, fleeting N-heterocyclic carbene (NHC)-supported $\{BH_2\}^-$ parent boryl anion (**II**) at low temperature and trapping it with a variety of electrophiles,^[3] while our own group isolated a nucleophilic borolyl anion that displayed not only classical salt metathesis but also single-electron transfer reactivity.^[4]

A major breakthrough in low-valent boron chemistry came with the isolation by Bertrand and coworkers of the first metal-free boron(I) compound, borylene **III**,^[5] which owes its remarkable stability to the excellent σ donor and π acceptor properties and the steric shielding afforded by the two supporting cyclic (alkyl)(amino)carbene (CAAC) ligands.^[6] Since then, CAACs have been successfully employed to stabilize a wide range of ever more reactive borylenes^[7] and boryl anions.^[8] The most recent examples of these include the CAAC-CO-stabilized derivative **IV**, which under photolytic conditions liberates CO and generates a dicoordinate borylene synthon,^[9] the highly reactive chloro(hydro)boryl anion **V**, which was isolated as a potassium crown ether complex,^[10] and most recently, the N_2 activation compound **VI**, which may be viewed as a N_2 -bridged borylene dimer.^[11]

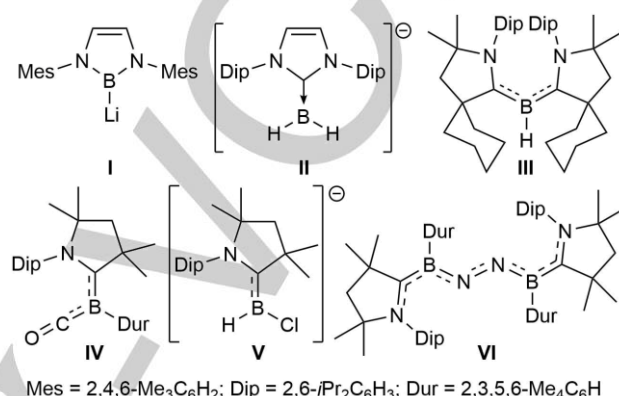
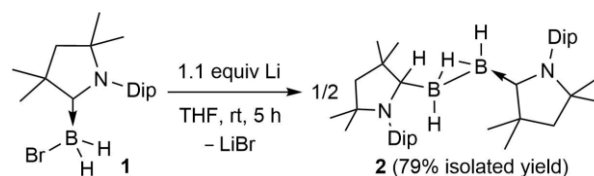


Figure 1. Selected examples of low-valent borylenes and boryl anions.

In this contribution we describe the facile stepwise reduction of a CAAC-stabilized (dihydro)haloborane to the corresponding diborane, and further to the first isolable, room-temperature-stable $\{BH_2\}^-$ parent boryl anion, which reacts as a strong boron nucleophile.

The room temperature reduction of $(CAAC^{Me})BH_2Br$ (**1**, see Fig. S21 in the SI for the solid-state structure of **1**) with 1.1 equiv. lithium sand in THF (Scheme 1) yielded a pale yellow solution exhibiting a single broad ^{11}B NMR resonance at 21.2 ppm (fwhm \approx 420 Hz). After removal of volatiles and extraction with hexanes, crystallization at $-25^\circ C$ yielded a crop of gold-orange crystals of compound **2** (79% yield). The $^1H\{^{11}B\}$ NMR spectrum of **2** showed a single, symmetrical $CAAC^{Me}$ ligand environment and a broad $2H\ B/H_2$ resonance at 2.66 ppm. The very broad ^{13}C NMR resonance of the carbene carbon atom was detected by an HMBC experiment at 161.5 ppm, 67 ppm upfield from that of the borane precursor **1** ($\delta_{13C} = 128.2$ ppm).



Scheme 1. One-electron reduction and homocoupling of **1** to form diborane **2**. Dip = 2,6-diisopropylphenyl.

The symmetry implied by the NMR data contrasts with the crystallographically-derived structure of **2**, which shows an

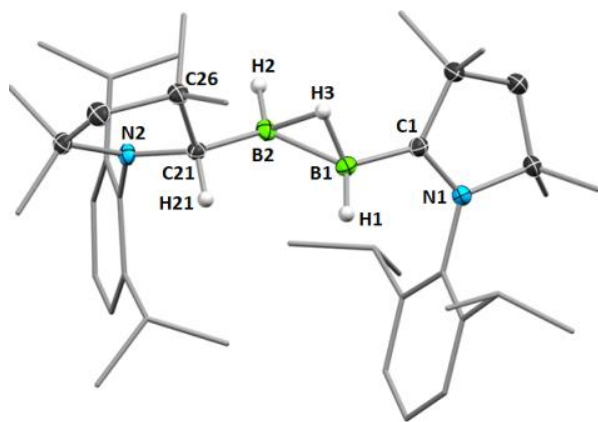
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unsymmetrically μ^2 -hydride-bridged diborane(5) (Fig. 2). B1 is supported by a neutral, purely σ -donating CAAC^{Me} ligand (B1-C1 1.5477(16) Å), whereas B2 is coordinated by a protonated CAAC^{Me} ligand (CAAC^{Me}H), which displays an sp^3 -hybridized C21 carbon atom (N2-C21-B2 114.15(9), C26-C21-B2 114.29(9)°) with B2-C21 and N2-C21 single bonds (1.6039(16) and 1.4774(14) Å, respectively). The B-B bond length of 1.6394(18) Å is significantly shorter than in related μ^2 -hydride-bridged diboranes(5) obtained from the reduction of mono-NHC-stabilized 1,1-diaryl-2,2-dichlorodiboranes (ca. 1.67 – 1.68 Å),^[12] presumably due to the absence of steric repulsion from the hydride ligands in **2**.

The synthesis of compound **2** is a rare example of targeted reduction of a neutral Lewis-base-stabilized sp^3 -borane to a neutral diborane. All previous examples involved NHC-stabilized di- and trihaloboranes undergoing both reductive coupling and exchange of all remaining halogens with hydrides, through hydrogen abstraction from the reaction solvent by radical intermediates.^[13] The formation of **2** may be facilitated by **1** undergoing a 1,2-hydride shift from boron to the CAAC^{Me} carbene carbon atom to form the isomeric sp^2 -borane (CAAC^{Me}H)BHB^r, **1'**, the free energy of which was calculated to be only 11.9 kcal·mol⁻¹ above that of **1** (see Fig. S24), thereby making it accessible under the reaction conditions employed in the reduction of **1** to **2**. Such reversible boron-to-carbon hydride shifts are well-documented in CAAC-supported hydroboron compounds^[15] and are owed to the good π acceptor properties of CAACs, which result in a relatively low-lying LUMO and a small HOMO-LUMO gap.^[6]



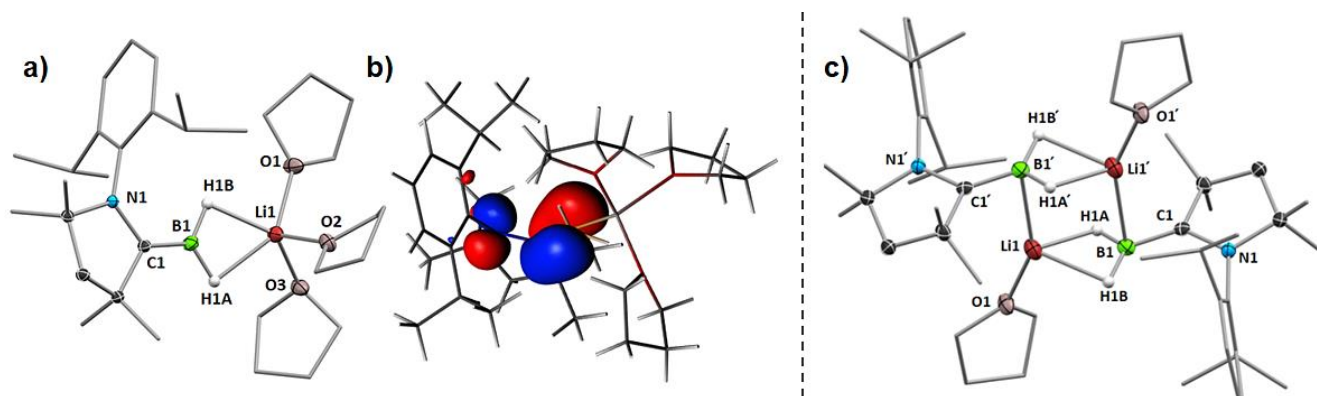
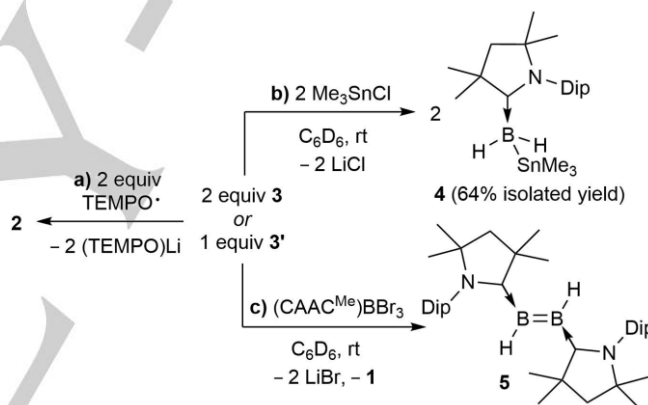


Figure 3. a) and c) Crystallographically-derived molecular structure of compounds **3** and **3'**, respectively. Thermal ellipsoids drawn at 50% probability level. Ellipsoids on the CAAC^{Me} ligand periphery and most hydrogen atoms omitted for clarity. Selected bond lengths (Å) and angles (°): **3** B1-C1 1.440(2), C1-N1 1.4332(15), B1-H1A 1.145(17), B1-H1B 1.140(16), B1...Li1 2.293(3), Li1-H1A 2.000(17), Li1-H1B 1.917(17), C1-B1-Li1 165.45(15), $\Sigma\angle_{B1}$ 356.0(9), $\Sigma\angle_{C1}$ 359.96(11); **3'** B1-C1 1.441(6), C1-N1 1.421(4), B1-H1A 1.11(4), B1-H1B 1.19(4), B1...Li1 2.301(8), B1...Li1' 2.407(8), Li1...Li1' 3.041(13), H1A-Li1 1.91(4), H1B-Li1 2.07(4), Li1...C1' 2.669(8), C1-B1-Li1' 162.9(4), C1-B1-Li 83.8(3), B1-Li1-B1' 99.6(3), $\Sigma\angle_{C1}$ 359.6(3), $\Sigma\angle_{B1}$ 360(2); b) Plot of the HOMO of **3** at the OLYP/TZ2P level of theory (−1.905 eV, isovalue 0.005).

structure of **3'** shows a boryl anion moiety with similar structural parameters to **3** (Fig. 3c). Its lithium cations are coordinated by a single THF residue and bridge over the boron-bound hydrides to the second boron center so as to form a distorted B₂Li₂ square (B1...Li1 2.301(8); B1...Li1' 2.407(8) Å; B1-Li1-B1' 99.6(3); Li1-B1-Li1' 80.4(3)°). It is noteworthy that the coordination of each Li cation to the boron atom of the second dimer moiety occurs perpendicularly, thereby allowing the CBH₂ borylene unit to remain planar. Unlike monomeric **3**, isolated crystals of dimeric **3'** proved virtually insoluble in hydrocarbon solvents, thus precluding the acquisition of NMR data. Though indefinitely stable in THF solution up to 70 °C under inert atmosphere, both **3** and **3'** slowly decomposed in the solid state to (CAAC)BH₃, even when stored in the glovebox freezer at −30 °C.

Plots of the frontier molecular orbitals (MOs) of **3** show a HOMO corresponding to the B-C π -bond, with a small π antibonding contribution from the C1-N1 bonds (Fig. 3b), similar to other CAAC-supported boryl anions.^[8,10] Furthermore, the B-C Mayer bond order amounts to 1.664, which is less than for the analogous [(CAAC^{Me})BHCl][−] anion (1.703),^[10] but still indicative of substantial π backbonding from boron to CAAC^{Me}. The calculated Hirshfeld partial charges of −0.161 for B1 and −0.077 for C1^[17] confirm the negative charge accumulation on the boron atom, which is the opposite charge distribution calculated for [(CAAC^{Me})BHCl][−] (−0.086 for B1 and −0.102 for C1).^[10] This should make **3/3'** particularly nucleophilic at boron.

Interestingly, cyclic voltammetry performed on compounds **1**, **2** and **3** in THF showed in all three cases a reversible reduction wave at −3.2 V (referenced to the (Fc⁺/Fc) couple, see Figs. S18 – S20), suggesting the possibility of a chemically reversible reduction of **2** to **3**. Indeed the reduction of isolated **2** with Li in THF led to the clean formation of **3**, a process that most likely ensues from the B-B bond cleavage of isomer **2'**. Conversely, the oxidation of **3/3'** with the 2,2,6,6-tetramethylpiperidyl radical (TEMPO) resulted in quantitative conversion back to diborane **2** (Scheme 4a), thus confirming the chemical reversibility of the reduction as well as of the **2/2'** hydride shift process.

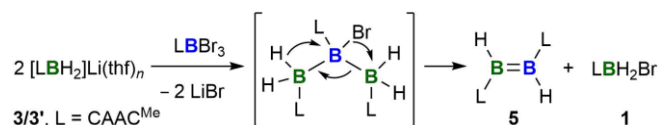


Scheme 4. Reactivity of **3/3'**.

The nucleophilic character of **3/3'** was confirmed by salt metathesis with Me₃SnCl, which quantitatively yielded colorless (CAAC^{Me})BH₂(SnMe₃) (**4**, Scheme 4b, see Fig. S23 for the solid-state structure of **4**), characterized by a ¹¹B NMR triplet at −29.7 (¹J_{B-H} = 100 Hz) and a broad ¹¹⁹Sn NMR quartet at −24.4 (¹J_{B-119Sn} = 314 Hz), effectively identical to that of the previously reported compound (CAAC^{Me})BH(CN)(SnMe₃).^[8a]

The reaction of **3** with one equiv. (CAAC^{Me})BBR₃ was accompanied by an instant color change to deep blue as well as the formation of a colorless precipitate, presumably LiBr. The ¹¹B NMR spectrum of the reaction mixture showed quantitative formation of **1** (broad triplet at δ_{11B} = −19.0, ¹J_{B-H} = 103 Hz) as well as a broad resonance at 40.0 ppm attributable to the blue-colored dihydrodiborene (CAAC^{Me})₂B₂H₂ (**5**, Scheme 4c),^[18] which was confirmed by high-resolution mass spectrometry ([C₄₀H₆₄B₂N₂]: calc. 594.5250; found 594.5227). We postulate that the reaction proceeds via a double salt metathesis leading to an intermediate triborane(7), which undergoes intramolecular bromide and hydride migration, as well as B-B bond cleavage, to generate **5** and **1** (Scheme 5). While this is certainly not the

most efficient way to synthesize diborene **5**, this unique reactivity has the potential to be harnessed for the synthesis of unsymmetrical B=B or new boron-element double bonds.



Scheme 5. Possible mechanism for the reaction of **3** with (CAAC^{Me})BBR₃.

In conclusion, we have shown that (CAAC^{Me})BH₂Br (**1**) undergoes a selective, B-B bond-forming, one-electron reduction to a hydride-shift isomer of the tetrahydridoborane (CAAC^{Me})₂B₂H₄, compound **2**. Subsequent two-electron reduction of **2** yields the first isolable parent boryl anion, [(CAAC^{Me})BH₂][−], which may be isolated in its monomeric form **3** or dimeric form **3'**, depending on the crystallization conditions. Compound **3** owes its remarkable solution stability to the strong π acceptor properties of the CAAC ligand, their HOMO being entirely delocalized over the B-C π bond. Furthermore, [(CAAC^{Me})BH₂][−] can be quantitatively oxidized back to **2**, in an unusual B-B bond-forming oxidation reaction with the TEMPO radical. While [(CAAC^{Me})BH₂][−] reacts as a typical boron nucleophile towards Me₃SnCl, leading to the formation of a new B-Sn bond, it undergoes a unique B=B double-bond-forming double salt metathesis with (CAAC^{Me})BBR₃, generating the dihydridoborene (CAAC^{Me})₂B₂H₂ and **1**. In view of these promising preliminary results, we are continuing to explore the reactivity of **3/3'** towards other p-block electrophiles.

Acknowledgements

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Conflict of interest

The authors declare no conflict of interest.

Keywords: boryl anion • diborane • reduction • nucleophilic boron • cyclic (alkyl)(amino)carbene

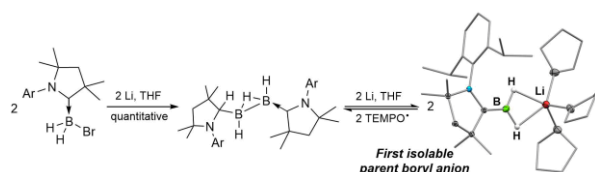
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Entry for the Table of Contents

Layout 2:

COMMUNICATION



M. Arrowsmith, J. D. Mattock, S. Hagspiel, I. Krummenacher, A. Vargas, H. Braunschweig*

Page No. – Page No.

Facile Synthesis of a Stable Dihydroboryl $\{BH_2\}^-$ Anion

Successive reductions of a cyclic (alkyl)(amino)carbene (CAAC)-supported (bromo)dihydroborane yield a hydride-shift isomer of the corresponding tetrahydrodiborane and the first stable dihydroboryl anion, which acts as a powerful boron nucleophile.